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Identifying mantle carbonatite metasomatism through Os–Sr–Mg isotopes in Tibetan ultrapotassic rocks



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ABSTRACT

Mantle-derived magmas at convergent plate boundaries provide unique insights into the nature of materials subducted to and recycled from depths. Here we present a study of Os-Sr-Mg isotopes on the Oligocene-Miocene ultrapotassic rocks aimed at better understanding sediment subduction and recycling beneath southern Tibet. New isotopic data confirm that ultrapotassic rocks in southern Tibet are of mantle origin, but underwent crustal contamination as evidenced by the variably high ¹⁸⁷Os/¹⁸⁸Os that obviously deviates from normal mantle reservoir. Still some samples with mantlelike 187 Os/ 188 Os exhibit δ^{26} Mg significantly lower than mantle and crustal lithologies, suggesting that the isotopically light Mg may not result from crustal contamination but retain specific fingerprint of carbonate-related metasomatism in mantle sources. Mantle carbonatite metasomatism is manifested by the inverse δ^{26} Mg $-^{87}$ Sr $/^{86}$ Sr correlations, as well as the depletion of high field strength elements relative to rare earth elements and the enrichment of CaO in ultrapotassic rocks. The positive co-variations between δ^{26} Mg and Hf/Sm defined by those low-¹⁸⁷Os/¹⁸⁸Os ultrapotassic rocks provide evidence for the potential of recycled dolomites to modify mantle Mg isotopic composition. The correlated spatial variations of δ^{26} Mg and Hf/Sm are interpreted to reflect carbonatitic metasomatism associated with the northward subduction of the Neo-Tethyan oceanic slab and its profound influence on postcollisional ultrapotassic magmatism.

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1. Introduction

Subduction of carbonate-bearing sediments at convergent plate boundaries has played a fundamental role in mantle refertilization (White and Patchett, 1984; Plank and Langmuir, 1998; Ducea et al., 2005) and in global carbon cycling (Connolly, 2005; Dasgupta and Hirschmann, 2010; Yang et al., 2012; Huang et al., 2015). Recently, volatile-rich mafic rocks including kimberlite, carbonatite, and potassic rocks of mantle origin (e.g. kamafugite and ultrapotassic rocks) have received an increasing amount of attention because of their petrogenetic links with carbonate-related metasomatism in deep earth (Thomsen and Schmidt, 2008; Avanzinelli et al., 2009; Liu et al., 2014b; Conticelli et al., 2015). Given the subduction of carbonate-rich Tethyan seafloor (cf. Johnston et al., 2011), the widespread mantle-derived ultrapotassic rocks in the Alpine–Himalaya orogenic belt thus offer a prime opportunity to investigate carbonatitic metasomatism and carbon recycling processes in subduction zone.

In southern Tibet, however, carbonatite metasomatism associated with the Neo-Tethyan seafloor subduction has long been understated (cf. Turner et al., 1996; Chung et al., 2005; Liu et al., 2014b), which will inhibit us to better understand the geochemical evolution of lithospheric mantle. Additionally, the inevitable crustal contamination, which has been amplified by crustal thickening in response to the India–Asia convergence (cf. Liu et al., 2014c), constitutes another difficulty in using mantle-derived rocks to probe the nature of mantle lithosphere. Selecting process-sensitive tracer

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Fig. 1. (a) Geological map of the Lhasa terrane, showing spatial distribution of the Mesozoic-Cenozoic magmatism and sampling locations in this study. (b) Geographic map of the Tibetan plateau. JSS = Jin Sha suture zone; BNS = Bangong-Nujiang suture zone; SNMZ = Shiquan river-Nam Tso Mélange zone; IYZS = Indus-Yarlung Zangbo suture zone; ATF = Altyn Tagh fault; LMF = Luobadui-Milashan fault; KF = Karakorum fault.

is therefore essential to ascertain the nature and extent of recycled seafloor materials with respect to the influence of crustal contamination.

Because Os behaves much more compatible than its radioactive parent element Re during mantle melting (Shirey and Walker, 1998), the residual mantle is strongly depleted in Re and exhibits Os concentration much higher than those of metasomatic components, which makes Re-Os isotopic system in mantle lithosphere essentially unaffected by subsequent metasomatic modification (Schaefer et al., 2000; Meisel et al., 2001). On the other hand, Os isotopic compositions of mantle-derived magmas are proved to be highly susceptible to crustal contamination due to low Os concentrations (Shirey and Walker, 1998; Schaefer et al., 2000), suggesting that Os isotopes in mantle-derived rocks can be utilized as a sensitive indicator for crustal contamination. Meanwhile, higher MgO contents in ultrapotassic rocks relative to crustal contaminants and limited Mg isotopic fractionation during high-temperature magmatic processes (Teng et al., 2010) enable Mg isotopic tracer to preserve signatures of mantle sources. The distinct Sr and Mg isotopic compositions between marine carbonates (Fantle and Higgins, 2014; Blättler et al., 2015) and terrestrial mantle reservoir (Teng et al., 2010) further indicate the potential of these isotopic proxies for mantle carbonatite metasomatism (Yang et al., 2012; Huang et al., 2015).

Here we present a systematic study on the Os, Sr, and Mg isotopes of the Tibetan ultrapotassic rocks. The combined application of multiple isotopes and trace element allow us to reveal the close relationship between the ultrapotassic rocks and mantle carbonatite metasomatism, and to better understand mantle refertilization processes during the northward subduction of Neo-Tethyan seafloor. As a case study, it also underscores the efficacy of Mg isotopes in mantle-derived rocks to provide information about carbon recycling in deep earth.

2. Geological setting

As a micro-continent rifted away from the eastern margin of Gondwana (cf. Yin and Harrison, 2000; Zhu et al., 2013), the Lhasa terrane in southern Tibet underwent a series of seafloor subduction episodes that have modified its mantle lithosphere (cf. Kapp et al., 2007; Ji et al., 2009; Zhu et al., 2013). As the northern boundary, the Bangong-Nujiang suture zone (BNS) separates Lhasa terrane from Qiangtang terrane (Fig. 1). To the south, the Indus-Yarlung Zangbo suture zone (IYZS) is viewed as remnants of the Neo-Tethyan seafloor lithologies between Lhasa terrane and India plate (cf. Yin and Harrison, 2000; Zhu et al., 2013). Recent studies reveal that the Lhasa terrane is composed of a reworked ancient crust in the center and juvenile additions to its northern and southern edges (cf. li et al., 2009; Zhu et al., 2013). Accordingly, the Lhasa terrane can be subdivided into the northern, central, and southern Lhasa subterranes, which are separated by the Shiquan river-Nam Tso Mélange zone (SNMZ) and Luobadui-Milashan fault (LMF) from north to south (Fig. 1a).

As products of the Tethyan seafloor subduction and the India-Asia continental collision thereafter, the Mesozoic-early Cenozoic magmatic rocks are distributed roughly parallel to the E–W trending Lhasa terrane (Fig. 1b). The Cretaceous volcanic rocks (131–91 Ma, Zhu et al., 2013) are widespread in the northern and central Lhasa subterranes. The southern Lhasa subterrane is dominated by the early Paleogene Linzizong volcanic sequence (69–43 Ma, Lee et al., 2009), some of which extending into the eastern part of the central Lhasa subterrane (Fig. 1a). The Gangdese plutons outcrop widely in the Lhasa terrane and can be roughly divided into the south Gangdese batholith (205–41 Ma, Ji et al., 2009; Zhu et al., 2013) associated with the IYZS and the north Gangdese batholith (194–110 Ma, Zhu et al., 2013) distributing along the BNS (Fig. 1a). With respect to the temporal and spatial distribution, the Mesozoic–early Cenozoic magmatism in the Lhasa terrane has been interpreted as reflecting continued Neo-Tethyan seafloor subduction, including low-angle northward subduction of the Indus–Yarlung Zangbo Neo-Tethyan seafloor, slab rollback and subsequent break-off (cf. Kapp et al., 2007; Ji et al., 2009). Another attractive explanation is the bi-direction seafloor subduction model (cf. Zhu et al., 2013), which is represented by southward subduction of the Bangong–Nujiang Neo-Tethyan seafloor and northward subduction of the Indus–Yarlung Zangbo Neo-Tethyan seafloor.

3. Ultrapotassic volcanism and sample description

Following the early-Paleogene India-Asia continental collision (Yin and Harrison, 2000; Lee et al., 2009), postcollisional magmatism in southern Tibet occurred in response to the continued India-Asia convergence (Turner et al., 1996; Chung et al., 2005). Mantle-derived ultrapotassic volcanic rocks (UPVR) commonly occurred as lava flows in the central and southern Lhasa subterranes (Fig. 1a), forming a magmatic belt with eruptive ages varying from 24 Ma to 10 Ma (Liu et al., 2014c). The mantle origin of the UPVR is indicated by mantle xenoliths (Liu et al., 2014a), whereas the entrained crustal xenoliths of varying size (e.g. gabbro and granite) also point to the significance of crustal contamination in petrogenesis of the UPVR (Liu et al., 2014c). As ultrapotassic magmatism in southern Tibet is nearly coeval with the accelerated plateau uplift and the east-west extension (cf. Yin and Harrison, 2000; Liu et al., 2014b), geodynamic models, such as convective thinning of mantle lithosphere (Turner et al., 1996), removal or delamination of overthickened lithosphere (Chung et al., 2005; Zhao et al., 2009), and roll-back and break-off of the subducted Indian plate (Guo et al., 2013), have been invoked to account for the mantle melting events during postcollisional stage.

We have collected a total of 47 UPVR samples with good spatial coverage of ultrapotassic magmatism across the Lhasa terrane (Fig. 1a). Two crustal xenolith samples from the host UPVR samples are also included in this study (Table 1). From basaltic trachyandesite to trachyte, this suite of UPVR exhibits a wide lithological range with phenocrysts composed of phlogopite, plagioclase, pyroxene, and minor olivine (Turner et al., 1996; Zhao et al., 2009). The compositional and mineralogical ranges of the studied UPVR are comparable to the full range of ultrapotassic rocks in southern Tibet (cf. review by Chung et al., 2005). High-quality zircon U–Pb age data for these UPVR samples have been reported (cf. Liu et al., 2014c), yielding the Oligocene–Miocene eruptive ages (24–10 Ma) that agree within error with K–Ar and Ar–Ar age data (Chung et al., 2005; Zhao et al., 2009). Sample details are listed in supplementary Table S1.

4. Analytical methods

In order to minimize the influence of chemical weathering on isotopic composition, weathered rock surfaces have been removed and thoroughly cleaned before analysis. After extracting crustal and mantle xenoliths by handpicking, fresh samples (the chemical index of alteration ranges from 27.6 to 51.0, Table S2) were powdered into 200 mesh. Major and trace element compositions are reported in Table S2. Analytical results of Re–Os, Sr, and Mg isotopes are given in Table 1.

4.1. Major and trace element compositions

Major and trace element analyses were conducted in the State Key Laboratory of Geological Processes and Mineral Resources (GPMR), China University of Geosciences, Wuhan. Rock powder (\sim 0.5 g) was mixed with 5.0 g compound flux (Li₂B₄O₇:LiBO₂ = 12:22) and fused in a Pt-Au crucible by heating at \sim 1050 °C

for 11 min. The mixture was swirled repeatedly to ensure complete molten and homogenization before pouring into a mold to form a flat disc for analysis. Major elements were analyzed using a *Shimadzu XRF-1800* sequential X-ray fluorescence spectrometry. The precision and accuracy for major element analysis are better than 4% and 3%, respectively.

For trace element analysis, the rock powder (\sim 50 mg) was dissolved in a Teflon bomb with HF + HNO₃ mixture and then was heated at 190 °C for 48 h. After evaporating the solution to dryness, the dried sample was re-dissolved using \sim 3 ml of 30% HNO₃, and then heated at 190 °C for 24 h. The final solution was diluted to \sim 100 g with 2% HNO₃ for subsequent analysis. Trace elements were analyzed using an *Agilent 7500a* inductively coupled plasma-mass spectrometry (ICP-MS). Analytical results of procedural blanks and international rock standards are given in Table S3.

4.2. Rhenium-osmium and strontium isotopic compositions

Re and Os isotopic analyses were conducted at the Guangzhou Institute of Geochemistry, Chinese Academy of Sciences (GIG-CAS), with some samples analyzed at the Miami University (Miami U.). In GIG-CAS, rock powders (~ 2 g), spiked with ¹⁸⁵Re and ¹⁹⁰Os, were frozen with inverse aqua regia into Carius tubes before being sealed and heated at 240 °C for 24 h. The separation of Re and Os was performed using a carbon tetrachloride aqua (CCl_4) solvent extraction method and Os was further purified by microdistillation. Purified Os was loaded using Ba(NO₃)₂ as emission enhancer on Pt filaments and determined using a Finnigan Triton thermal ionization mass spectrometer (TIMS) operated in the negative ion mode. Re was separated and purified from the liquid remaining after extraction of Os using AG50W-X8 (200-400 mesh) cation exchange columns and BPHA (100 mesh) extraction chromatographic columns. Isotope dilution analysis of Re was conducted on a Thermo-Scientific X series-2 ICP-MS. Instrumental mass fractionation of Os was corrected by normalizing the measured ¹⁹²Os/¹⁸⁸Os to 3.08271. Analytical uncertainty for Os concentration and ¹⁸⁷Os/¹⁸⁸Os are less than 0.5% within two times of the standard deviation (2σ) . Total procedural blanks for Re and Os were \sim 20 pg and 0.18 pg, respectively, yielding average ¹⁸⁷Os/¹⁸⁸Os of 0.350 (Table S4). The contribution of the blank to measured Os concentration and ¹⁸⁷Os/¹⁸⁸Os were <3% and <10%, respectively.

In the Miami U., rock powders (~ 1 g) were spiked with ¹⁸⁵Re and ¹⁹⁰Os and digested with concentrated inverse aqua regia in sealed Carius tubes by heating at 230 °C for 48 h. The separation of Re and Os was performed using a CCl₄ solvent extraction method, and Os was further purified by microdistillation. Re was separated and purified from the liquid remaining after extraction of Os using AG1-X8 anion exchange columns. Purified Os was loaded on Pt filaments along with Ba(OH)₂ as emission enhancer, and Os isotope ratio was determined using negative TIMS as OsO₃⁻⁻ using *Finnigan Triton* TIMS. Os isotope ratios were corrected for mass fractionation using ¹⁹²Os/¹⁸⁸Os = 3.0826. Re concentrations were measured by isotope dilution by negative TIMS as ReO₄⁻⁻. Total procedural blanks were ~2 pg for Re and ~0.1 pg for Os. The average ¹⁸⁷Os/¹⁸⁸Os of unspiked blanks was 0.184 (Table S4).

The Sr isotopic analyses were carried out on a *Micromass ISO-PROBE* MC-ICP-MS at GIG-CAS. Rock powders (50–100 mg) were dissolved in screw-top PFA beakers using a 1:2 (v/v) mixture of HNO₃ and HF acids. The capped beakers were heated at 140 °C for 7 days before opened and then evaporated to dryness. The separation of Sr was done using cation exchange columns. All the measured Sr isotopic ratios were corrected for mass fractionation using 86 Sr = 0.1194. Multiple analyses of the BHVO-2 and NBS-987 yield average 87 Sr/ 86 Sr of 0.703469 ± 0.000010 (2 σ , *n* = 2) and 0.710268 ± 0.000010 (2 σ , *n* = 7), respectively (Table S5).

Table 1 Os–Sr–Mg isotopic compositions of Tibetan ultrapotassic rocks and crustal xenoliths.

Sample No.	Locality	Lith.	Mg [#]	Hf/Sm	Re	Os	¹⁸⁷ Re/ ¹⁸⁸ Os	¹⁸⁷ Os/ ¹⁸⁸ O)s	⁸⁷ Sr/ ⁸⁶ Sr		δ^{25} Mg	$\pm 2\sigma$	$\delta^{26} { m Mg}$	$\pm 2\sigma$
					(ppb)	(ppb)		Ratio	$\pm 2\sigma$	Ratio	$\pm 2\sigma$	(‰)		(‰)	
Ultrapotassic vo	lcanic rocks (U	PVR)													
CQ01*	Maiga	BtA	76.0	0.38	0.047	0.199	1.13	0.1324	0.0001	0.722810		-0.19	0.09	-0.28	0.08
CQ02*	Maiga	TrA	56.5	0.66	0.051	0.076	3.23	0.1509	0.0001	0.716194		-0.13	0.09	-0.26	0.08
CQ03*	Maiga	BtA	73.5	0.38	0.016	0.089	0.87	0.1441	0.0001	0.724697		-0.19	0.09	-0.35	0.08
D9103*	Maiga	TrA	61.6	0.66	0.044	0.081	2.63	0.1572	0.0002	0.719414		-0.19	0.09	-0.40	0.08
GGP-7*	Maiga	Te	75.6	0.27	0.070	0.128	2.65	0.1423	0.0001	0.715817		-0.20	0.09	-0.35	0.08
Os-Rep	Maiga	Те			0.093	0.087	5.18	0.1492	0.0001						
XR01-3*	Chazi	BtA	75.7	0.38	0.073	0.020	17.9	0.2226	0.0003	0.721161		-0.09	0.09	-0.29	0.08
Mg-Rep	Chazi	BtA										-0.13	0.05	-0.30	0.06
XR02-1*	Chazi	TeP	67.7	0.52	0.014	0.698	0.09	0.1315	0.0001	0.736451		-0.18	0.10	-0.30	0.05
XR1102	Chazi	TrA	64.9	0.60	0.004	0.015	1.14	0.1924	0.0008	0.716657	0.000011	-0.15	0.03	-0.29	0.01
DR01-2*	Mibale	lr.	57.9	0.96	0.056	0.016	17.2	0.1710	0.0002	0.718043		-0.27	0.10	-0.42	0.05
28030-18*	Tangra	TrA	61.4	0.68	0.055	0.069	3.87	0.1647	0.0001	0.719632					
Os-Rep	YumCo	IrA	64.9	0.74	0.058	0.097	2.90	0.1567	0.0001	0 5105 45	0.000040	0.00	0.01	0.44	0.00
DR1103	Mibale	IrA TrA	64.3	0.71	0.075	0.066	5.58	0.1904	0.0003	0./19/45	0.000012	-0.22	0.01	-0.44	0.03
DRIIU/	Mibale	TrA TrA	C2 3	0.68	0.088	0.056	7.03	0.1746	0.0004	0.718702	0.000013	-0.20	0.01	-0.36	0.09
DRIII3	Mibale	TrA TrA	51.0	0.71	0.019	0.045	2.06	0.1734	0.0005	0.719773	0.000012	-0.24	0.07	-0.45	0.06
DK1114 7D1*	Zabuwa	TrA TrA	51.0	0.67	0.022	0.045	2.30	0.1934	0.0005	0.719901	0.000011	-0.25	0.04	-0.46	0.04
ZD1 7D4*	Zabuye	TrA TrA	50.1	0.50						0.711352		-0.11	0.04	-0.26	0.05
ZD4 7D10*	Zabuye	TrA	57.0	0.50						0.711075		-0.11	0.04	-0.27	0.05
ZD12 7P1102	Zabuye	TrA	55.2	0.60	0 122	0.020	21.6	0 22/1	0.0005	0.712408	0.000011	-0.12	0.04	-0.20	0.03
ZD1102 7B1108	Zabuye	TrA	56.1	0.87	0.132	0.030	21.0	0.2341	0.0005	0.711252	0.000011	-0.13	0.02	-0.28	0.02
ZB1100 7B1112	Zabuye	Tr A	55.7	0.70	0.121	0.028	21.2	0.2288	0.0005	0.711275	0.000011	-0.12	0.05	-0.22	0.03
SL0621*	Sailinu	TrA	72 A	0.07	0.105	0.130	3 51	0.1747	0.0003	0.711241	0.000010	-0.18	0.07	-0.30	0.05
Os-Ren	Sailipu	TrA	12.4	0.45	0.054	0.130	3 51	0 1684	0.0001			0.10	0.07	0.27	0.00
SL0626*	Sailipu	TrA	64 9	0.53	0.025	0.105	1 15	0 1624	0.0002						
SL0630*	Sailipu	TrA	73 7	0.42	0.026	0.010	12.1	0 3067	0.0004			-0.18	0.10	-0.30	0.05
SLP1101	Sailipu	TrA	71.1	0.51	0.065	0.128	2.48	0.1707	0.0003	0.719810	0.000013	-0.17	0.08	-0.31	0.05
XN1201	Xuena	TrA	61.8	0.50	0.024	0.009	13.0	0.5956	0.0016	0.711178	0.000012	-0.15	0.03	-0.29	0.03
XN1203	Xuena	TrA	61.4	0.52	0.031	0.009	16.6	0.6191	0.0023	0.711569	0.000010	-0.13	0.03	-0.24	0.05
XN1206	Xuena	TrA	64.7	0.50	0.005	0.016	1.72	0.3653	0.0012	0.711474	0.000012	-0.15	0.06	-0.28	0.02
XN1207	Xuena	TrA	61.1	0.56	0.016	0.011	7.29	0.4717	0.0012	0.710747	0.000011	-0.14	0.01	-0.25	0.04
XN1208	Xuena	TrA	65.6	0.51	0.025	0.007	18.6	0.5081	0.0022	0.713008	0.000012	-0.17	0.02	-0.31	0.03
10XB11	XB-YR	TrA	65.6	0.64	0.006	0.040	0.71	0.1596	0.0004						
10XB15	XB-YR	Tr	61.0	0.61	0.013	0.096	0.64	0.1677	0.0003						
10YR02	XB-YR	TrA	61.7	0.58	0.416	0.035	58.8	0.3461	0.0009						
10YR06	XB-YR	TrA	69.7	0.57	0.044	0.022	9.66	0.3299	0.0009						
XB1207	XB-YR	TrA	66.2	1.00	0.102	0.014	35.7	0.2644	0.0009	0.717597	0.000012	-0.14	0.09	-0.27	0.05
XB1210	XB-YR	TrA	69.5	0.30	0.083	0.004	109.6	0.7197	0.0034	0.720108	0.000010	-0.19	0.07	-0.35	0.02
XB1214	XB-YR	TrA	67.3	0.36	0.052	0.004	58.6	0.3466	0.0015	0.718906	0.000012	-0.14	0.01	-0.30	0.02
XB1230	XB-YR	TrA	61.5	0.59	0.037	0.043	4.19	0.1736	0.0004	0.727526	0.000012	-0.18	0.03	-0.34	0.02
XB1232	XB-YR	Tr	52.8	0.72	0.033	0.095	1.68	0.1863	0.0004	0.724916	0.000013	-0.17	0.00	-0.33	0.03
XB1233	XB-YR	TrA	62.5	0.55	0.055	0.037	7.25	0.1661	0.0004	0.722197	0.000011	-0.17	0.08	-0.30	0.07
XB1234	XB-YR	TrA	63.4	0.61	0.057	0.034	8.18	0.1773	0.0006	0.723080	0.000011	-0.14	0.01	-0.28	0.04
XB1237	XB-YR	TrA	64.3	0.73	0.096	0.055	8.41	0.1639	0.0004	0.723372	0.000011	-0.15	0.04	-0.31	0.05
XB1241	XB-YR	TrA	63.4	0.47	0.069	0.054	6.18	0.2024	0.0004	0.724377	0.000011	-0.17	0.03	-0.36	0.05
XB1242	XB-YR	TrA	65.9	0.47	0.065	0.008	43.9	0.5659	0.0026	0.725390	0.000012	-0.17	0.02	-0.34	0.07
XB1253	XB-YR	TrA	69.1	0.66	0.039	0.016	11.8	0.3088	0.0007	0.719685	0.000011	-0.13	0.05	-0.27	0.02
XB1254	XB-YK	IrA T=A	69.I	0.65	0.119	0.023	25.5	0.3322	0.0006	0.719629	0.000011	-0.15	0.03	-0.28	0.08
AB1257		II'A D+A	71.9	0.34	0.019	0.050	1.89	0.2218	0.0006	0.713924	0.000010	-0.10	0.02	-0.32	0.04
YR1728	ХВ-ҮК	BtA	/1.4	0.34	0.008	0.054	0.69	0.1953	0.0005	0./138//	0.000011	-0.12	0.03	-0.29	0.04

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Sample No.	Locality	Lith.	$Mg^{\#}$	Hf/Sm	Re	Os	¹⁸⁷ Re/ ¹⁸⁸ Os	¹⁸⁷ Os/ ¹⁸⁸ O	łs	⁸⁷ Sr/ ⁸⁶ Sr		δ ²⁵ Mg	$\pm 2\sigma$	δ ²⁶ Mg	$\pm 2\sigma$
					(ddd)	(qdd)		Ratio	$\pm 2\sigma$	Ratio	$\pm 2\sigma$	(%)		(‰)	
Crustal xenoliths															
XB1260	XB-YR	Gabbro	55.8	0.52	0.132	0.034	18.79	0.2249	0.0007	0.709763	0.000013	-0.07	0.01	-0.12	0.06
XB1261	XB-YR	Granite	47.6	0.91						0.713443	0.000011	-0.24	0.04	-0.49	0.05
* Elemental and S	r isotopic con	npositions of	these samp	les have beei	ר reported in Zh	ao et al. (2009).	Their Os and Mg	isotopes were	: determined i	n the Miami Ur	niversity and th	e University of	Arkansas, re:	spectively.	
Os-Rep = replicate	Os data for	separate dige.	stions of di	fferent sampi	e aliquots. Mg-R	<i>lep</i> = repeat sar	nple dissolution, co	olumn chemis	stry and instru	imental analysis					
$2\sigma = 2$ times the	standard devi	iation of the I	population 4	of four repea	t measurements	of a sample so	lution.								
XB-YR, Xungba ar	d Yare area;	BtA, basaltic 1	trachyandes	ite; Te, tephr	ite; TeP, tephrip	honolite; Tr, tra	chyte; TrA, trachya	ndesite.							
Samples XB1257	nd XB1258 a	ire the host u	ltrapotassic	rocks for gal	broic (XB1260)	and granitic (XI	31261) xenoliths, r	espectively.							

Table 1 (continued)

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4.3. Magnesium isotope compositions

The Mg isotopic analyses, using the sample-standard bracketing method, were conducted using a Nu Plasma HR MC-ICP-MS at the Isotope Laboratory of the University of Arkansas and using a Neptune plus MC-ICP-MS at the China University of Geosciences, Beijing, respectively. Rock powders (1-20 mg) were dissolved in screw-top beakers using a 3:1 (v/v) mixture of Optimagrade HF and HNO₃ acids. These capped beakers were heated at 70-80°C, and then the solutions were evaporated to dryness. The dried samples were refluxed with a 3:1 (v/v) mixture of HCl and HNO₃, and then evaporated again to achieve 100% dissolution. The residues were then dissolved in 1N HNO3 for chromatographic separation. Separation of Mg was achieved using cation exchange columns loaded with 1 ml pre-cleaned resin (BioRad 200-400 mesh AG50W-X8), with 1N HNO₃ used as leaching agent (Teng et al., 2010). The Mg yields range from 99.5% to 99.9%. Samples were passed through the columns twice to achieve the required level of purification. Total procedural blank for Mg was <15 ng. The analytical results for magnesium isotopes are expressed as δ -notation in per mil relative to DSM3: $\delta^{X}Mg = [({}^{X}Mg/{}^{24}Mg)_{sample}/({}^{X}Mg/{}^{24}Mg)_{DSM3} - 1] \times 1000$, where X refers to mass 25 or 26 and DSM3 is a magnesium solution made from pure Mg metal (Galy et al., 2003). The analyses of AGV-2, BCR-2, BHVO-2, Kilbourne Hole olivine, and Hawaii seawater yield weighted average δ^{26} Mg values in good agreement with recommended values within 2σ error (Table S6).

5. Results

The UPVR samples have broad ranges of SiO₂ (45.4–60.6 wt.%), MgO (2.9–12.3 wt.%), CaO (1.9–10.1 wt.%), and K₂O (5.0–8.3 wt.%) (Table S2). Most UPVR samples are basaltic trachyandesite and trachyandesite, with a few samples plotted in the fields of tephrite (GGP-7), tephriphonolite (XR02-1), and trachyte (DR01-2, 10XB15, and XB1232) (Fig. 2a). Both gabbroic (SiO₂ = 50.1 wt.%) and granitic (SiO₂ = 69.7 wt.%) xenoliths have been identified from ultrapotassic rocks (Fig. 2a). In the primitive mantle-normalized incompatible trace element diagram (Fig. 2b), a remarkable feature of the UPVR is the depletion of high field strength elements (HFSEs) relative to rare earth elements (REEs). Negative anomalies in Zr and Hf can be found in the UPVR (Fig. 2b), and Hf/Sm ratio decreases with increasing Mg[#] and CaO/Al₂O₃ (Figs. 2c and 2d). Unlike the UPVR, Zr and Hf anomalies are rather weak for gabbroic and granitic xenoliths (Fig. 2b).

The UPVR samples display wide ranges of Re (0.004–0.416 ppb) and Os (0.004–0.698 ppb), and have more radiogenic Os isotopic compositions ($^{187}Os/^{188}Os = 0.1315-0.7197$, Table 1) than the primitive upper mantle ($^{187}Os/^{188}Os = 0.1296$, Meisel et al., 2001) and the entrained peridotite xenoliths ($^{187}Os/^{188}Os = 0.1246$, Table S7). Os concentration and $^{187}Os/^{188}Os$ ratio in gabbroic xenolith are 0.034 ppb and 0.2249, respectively. In the plot of $^{187}Os/^{188}Os$ vs. 1/Os (Fig. 3a), a scattered but positive correlation can be observed in the UPVR samples. The Tibetan UPVR samples also exhibit extremely radiogenic Sr isotopic compositions ($^{87}Sr/^{86}Sr = 0.710747-0.736451$, Table 1) as discussed previously (Chung et al., 2005; Zhao et al., 2009). The relatively low $^{87}Sr/^{86}Sr$ in gabbroic (0.709763 ± 0.000013) and granitic xenoliths (0.713443 ± 0.000011) are distinguishable from the host UPVR samples.

Compared with mantle peridotite (δ^{26} Mg = $-0.25 \pm 0.04\%$, Teng et al., 2010), the δ^{26} Mg values of mantle-derived ultrapotassic rocks are more variable (-0.22 to -0.46%, Table 1). Heavy Mg isotopic composition shown by gabbroic xenolith (δ^{26} Mg = $-0.12 \pm 0.06\%$) is comparable with that of lower continental crust (average δ^{26} Mg = -0.19%, Teng et al., 2013). In contrast, granitic xenolith has a light Mg isotopic composition, with δ^{26} Mg



Fig. 2. (a) Total alkalis vs. SiO₂ diagram, with alkaline/sub-alkaline dividing line. (b) Primitive mantle-normalized incompatible element patterns for the UPVR and the entrained crustal xenoliths. Primitive mantle data are from Sun and McDonough (1989). Dashed lines with different styles represent the incompatible element compositions of calcite (Ducea et al., 2005) and dolomite (Sapienza et al., 2009) in mantle peridotites. (c) Hf/Sm vs. Mg[#], and (d) Hf/Sm vs. CaO/Al₂O₃, showing that primitive ultrapotassic rocks (high Mg[#]) have higher CaO/Al₂O₃ (i.e., high CaO content) and are more depleted in HFSEs (i.e., Hf/Sm). Major and trace element data for the Tibetan UPVR (cf. Zhao et al., 2009; Liu et al., 2014b) and clinopyroxenes from mantle xenoliths (see references in Fig. S1) are plotted for comparison.

as low as $-0.49 \pm 0.05\%$ (Table 1). In the plot of δ^{25} Mg vs. δ^{26} Mg (Fig. 3b), the measured Mg isotopes follow the terrestrial massdependent equilibrium fractionation trend with a slope close to 0.521. According to the δ^{26} Mg $^{-187}$ Os 188 Os correlation (Fig. 3c), two kinds of UPVR samples can be identified: (i) samples with varying 187 Os 188 Os and mantle-like δ^{26} Mg (trend A), and (ii) samples showing restricted 187 Os 188 Os but variably lower δ^{26} Mg (trend B).

6. Discussion

6.1. Crustal contamination effects on Os and Mg isotopic compositions of ultrapotassic crocks

Tibetan ultrapotassic rocks have long been treated as primitive melts of mantle peridotite (e.g. Zhao et al., 2009; Guo et al., 2013) due to the entrainment of mantle xenoliths (Liu et al., 2014a) and high $Mg^{\#}$ (up to 76) and compatible element (e.g. Cr, Co, Ni) concentrations. However, caution is required in using isotope data of mantle-derived rocks to infer the nature of subcontinental lithospheric mantle (SCLM) because of the likely crustal contamination/assimilation in orogenic belts (Schaefer et al., 2000; Liu et al., 2014b). In southern Tibet, entrained crustal fragments and zircon xenocrysts of crustal origin indicate that both upper and lower crustal materials of the Lhasa terrane have been incorporated into ultrapotassic magma (Liu et al., 2014c). For this reason, the effects of crustal contamination on isotopic compositions of ultrapotassic rocks should be assessed before using these proxies to reveal mantle geochemical evolution beneath southern Tibet.

6.1.1. The susceptibility of Os isotopes to crustal contamination

During crust-mantle segregation and mantle partial melting, Re behaves as a moderately incompatible element while Os is strongly compatible (Shirey and Walker, 1998), which results in high Os concentration and uniform ¹⁸⁷Os/¹⁸⁸Os in residual mantle (Meisel et al., 2001). Such partitioning behaviors also lead to high Re/Os in continental crust that consequently develop variably high ¹⁸⁷Os/¹⁸⁸Os through radiogenic ingrowth of ¹⁸⁷Os from the β -decay of ¹⁸⁷Re (Saal et al., 1998; Peucker-Ehrenbrink and Jahn, 2001). The upper continental crust is characterized by low Os concentration and extremely high 187 Os/ 188 Os (e.g. Os = 0.031 ppb, 187 Os/ 188 Os = 1.4 ± 0.3, Peucker-Ehrenbrink and Jahn, 2001). Compared with upper continental crust, the lower continental crust has higher Os concentration and less radiogenic Os isotopic composition (average Os = 0.109 ppb, 187 Os/ 188 Os = 0.795, Saal et al., 1998). Because the Lhasa terrane has undergone episodes of juvenile crust formation in the Mesozoic-Cenozoic (cf. Ji et al., 2009; Zhu et al., 2013), the Os isotopic composition of lower crust would be more heterogeneous and unradiogenic because the juvenile crust additions (e.g. gabbroic xenoliths, sample XB1260) commonly yield low ¹⁸⁷Os/¹⁸⁸Os as a result of short crust formation age and limited radiogenic ingrowth of ¹⁸⁷Os.

As illustrated in Fig. 3a, simple magma-mixing process cannot account for the UPVR samples with high ¹⁸⁷Os/¹⁸⁸Os unless a large amount of crustal materials are incorporated into primitive ultrapotassic magma. Together with the convex upward $\delta^{18}O^{-87}$ Sr/⁸⁶Sr correlations (i.e., the crustal contamination trend defined by O and Sr isotopes, Liu et al., 2014b), the presence of crustal xenoliths and zircon xenocrysts in the UPVR (Liu et al., 2014c) demonstrates the occurrence of crustal assimilation during magmatic evolution. Furthermore, the fractional crystallization of olivine and sulfide will dramatically decrease Os concentrations of remaining magmas



Fig. 3. Os and Mg isotopic compositions of the Tibetan UPVR and entrained crustal xenoliths. (a) $^{187}Os/^{188}Os$ vs. 1/Os. Os isotopic compositions for three presumed crustal contaminants are given bellow: contaminant 1 (Os = 0.03 ppb, $^{187}Os/^{188}Os = 1.4$), contaminant 2 (Os = 0.045 ppb, $^{187}Os/^{188}Os = 0.85$), and contaminant 3 (Os = 0.1 ppb, $^{187}Os/^{188}Os = 0.25$). *R* is the ratio between crustal assimilation rate and fractional crystallization rate, and D_{Os} is the bulk partition coefficient for Os. The labeled percentages along modeling curves donate the AFC increments. Os isotopic variation predicted by magma mixing model is shown for comparison. Symbol sizes are greater than the 10% analytic uncertainties. (b) $\delta^{25}Mg_{DSM3}$, showing the Mg isotopic data in this study is consistent with the mass-dependent fractionation trend. Error bars of Mg isotopic ratios represent two standard deviations, and β represents the exponent in the mass-dependent fractionation law (Young and Galy, 2004). (c) $\delta^{26}Mg_{DSM3}$, showing to so and Mg isotopes of the UPVR samples and gabbroic xenolith. Mantle carbonatite metasomatism is modeled in terms of binary mixing between carbonate metls (dolomite and calcite) and mantle lherzolite. Os and Mg isotopic compositions of end-members are given in Table S7. Mixing hyperbolas are marked in 10% increments. Based on Os isotopic variation in Fig. 3a, R = 0.8 and $D_{Os} = 10$ are used in modeling AFC processes. Assuming cumulates of phlogopite + clinopyroxene + olivine, bulk partition coefficient for Mg (D_{Mg}) is set at 4 ± 2 . The uncertainty in D_{Mg} is indicated by color-shaded area with dashed lines for each solid AFC modeling curve. Circles on the AFC modeling curves indicate 2% AFC increments. Os and Mg isotopic ranges of normal mantle reservoir (Meisel et al., 2001; Teng et al., 2010) are shown for comparison.

(Saal et al., 1998), and result in their Re-Os isotope system being extremely susceptible to crustal contamination. Based on aforementioned Os isotopic signatures of continental crust, we modeled the assimilation and fractional crystallization (AFC) processes and found that AFC modeling can fully reproduce Os isotopic variation observed in the Tibetan ultrapotassic rocks (Fig. 3a). For the UPVR samples with high ¹⁸⁷Os/¹⁸⁸Os (>0.3), their radiogenic Os isotopic compositions may imply the extensive crustal contamination with large amounts of high-¹⁸⁷Os/¹⁸⁸Os crustal materials (e.g. upper crust and lower crust with sedimentary origin) incorporated into ultrapotassic magma (Fig. 3a). The UPVR samples with ¹⁸⁷Os/¹⁸⁸Os slightly deviated from the mantle value reflect magmas undergoing low-level crustal contamination and assimilating crustal materials showing ¹⁸⁷Os/¹⁸⁸Os as low as the entrained gabbroic xenolith (Fig. 3a). Therefore, considering the Oligocene-Miocene age of ultrapotassic magmatism (24-10 Ma, Liu et al., 2014c), the radiogenic Os isotopic compositions corroborate the role of crustal contamination/assimilation in petrogenesis of the Tibetan ultrapotassic rocks.

6.1.2. Limited effects of crustal contamination on Mg isotopes of the UPVR

But, unlike the Re–Os isotopic system, crustal contamination cannot fully account for the overall light Mg isotopic signatures

in the Tibetan ultrapotassic rocks (Fig. 3c), especially for the low-187Os/188Os UPVR samples that exhibit light Mg isotopic compositions (trend B in Fig. 3c). Recent studies reveal that light Mg isotope is prone to partition into liquid during chemical weathering and heavy Mg isotope will be left in the residue (cf. Li et al., 2010). Due to the influence of surface weathering, the average Mg isotopic compositions of upper (δ^{26} Mg = -0.22%, Li et al., 2010) and lower continental crust ($\delta^{26}Mg = -0.18\%$, Teng et al., 2013) are proved to be slightly heavier than mantle peridotites $(\delta^{26}Mg = -0.25 \pm 0.04\%)$ and basalts (MORB: $\delta^{26}Mg = -0.25 \pm$ 0.06%; OIB: $\delta^{26}Mg = -0.26 \pm 0.08\%$) (Teng et al., 2010). In this case, assimilating continental crust components like the entrained gabbroic xenolith (δ^{26} Mg = $-0.12 \pm 0.06\%$, Table 1) would only make Mg isotopes of ultrapotassic magma heavier than the mantle reservoir, contrasting with the overall light Mg isotopic compositions in the Tibetan UPVR (Fig. 3b).

Even if the heterogeneity in Mg isotopic composition of continental crust is considered (Li et al., 2010; Teng et al., 2013), higher MgO contents in mantle-derived ultrapotassic rocks than crustal contaminants further indicate that there is little leverage to change Mg isotopic compositions of ultrapotassic magmas through low-level crustal contamination. This suggestion is supported by the mantle-like Mg isotopic composition shown by the host UPVR sample (MgO = 8.09 wt.%, δ^{26} Mg = $-0.29 \pm 0.04\%$) of granitic



Fig. 4. Diagrams of (a) CaO/SiO₂ vs. MgO and (b) $\delta^{18}O_{V-SMOW}$ vs. SiO₂, showing isotopic and element variations of the Tibetan UPVR and the Italian ultrapotassic rocks that underwent contamination of the Apennine carbonates (cf. Peccerillo et al., 2010). Oxygen isotopic data of the Tibetan UVPR are from Zhao et al. (2009), Liu et al. (2014b) and references therein. Note that CaO contents of the Tibetan UPVR are apparently lower than those of the Italian ultrapotassic rocks that were contaminated by dolostone and limestone during magmatic evolution. The positive $\delta^{18}O_{V-SMOW}$ -SiO₂ correlation defined by the Tibetan UPVR contrasts with the negative trends found in the Italian ultrapotassic rocks, precluding contamination of sedimentary carbonates as a controlling process in generating isotopic signatures of the Tibetan UPVR.

xenolith (MgO = 0.61 wt.%, δ^{26} Mg = $-0.49 \pm 0.05\%$) (Table 1). Additionally, the correlations between Os and Mg isotopes provide evidence for the limited role of crustal contamination in changing Mg isotopic compositions of the UPVR (Fig. 3c). As predicted by AFC modeling (Fig. 3c), crustal contamination gives rise to dramatic increases in ¹⁸⁷Os/¹⁸⁸Os of mantle-derived ultrapotassic magmas (trend A in Fig. 3c). Meanwhile, substantial changes in δ^{26} Mg are not observed until large amounts of low-MgO crustal materials, such as pelites (MgO = 3.55 wt.%; δ^{26} Mg = $-0.52 \pm 0.07\%$, Li et al., 2010) and felsic granulite (MgO = 2.71 wt.%, δ^{26} Mg = -0.72 \pm 0.08^{\%}, Teng et al., 2013), are incorporated into ultrapotassic magmas. The contrasting effects of crustal contamination on Os and Mg isotopic variations can still be observed in AFC modeling even if the MgO content of primitive magma is assumed to be as low as 8 wt.%. We thus conclude that the overall light Mg isotopic signature, particularly for the variably low δ^{26} Mg in those less contaminated UPVR samples (87 Os/ 188 Os < 0.3), may not be result of low-level crustal contamination.

6.1.3. Contaminated by carbonate wall-rocks during magmatic ascent?

Unlike silicate crust components, sedimentary carbonate is an important low- δ^{26} Mg reservoir in the Earth (-5.28 to -1.09‰, Young and Galy, 2004), which means that incorporation of Mgrich carbonates (e.g. dolostone) can shift δ^{26} Mg of the UPVR to lower values. Moreover, previous works regarding the interaction between carbonate wall-rocks and ultrapotassic magmas have shown that incorporation of carbonates during magmatic evolution will result in significant increases of bulk-rock δ^{18} O_{V-SMOW} (up to 13.3‰) and CaO content (up to 23.05 wt.%, Peccerillo et al., 2010) (Fig. 4). Incorporation of CaO-rich mineral phases during magmatic evolution (e.g. CaO content in olivine is up to 1.63 wt.%, Wenzel et al., 2002).

Notwithstanding, many lines of evidence preclude the interaction between sedimentary carbonates and ultrapotassic magma as a controlling process to generate the Mg and O isotopic signatures observed in the Tibetan UPVR. First, carbonate xenoliths or globules have not been identified from the Tibetan ultrapotassic rocks so far (cf. Liu et al., 2014b), and phenocrysts such as olivines in the UPVR are reported to have low CaO contents (<0.25 wt.%, Turner et al., 1996; Zhao et al., 2009). In addition, the Tibetan UPVR samples display poorly varying and lower CaO/SiO₂ (<0.2, Fig. 4a), which apparently differs from the Alban Hills ultrapotassic rocks that underwent carbonate contamination (Peccerillo et al., 2010). Another argument against carbonate contamination comes from the positive correlation between $\delta^{18}O_{V-SMOW}$ and SiO₂ de-

fined by the Tibetan UPVR (Fig. 4b), which imply that the elevation of bulk-rock $\delta^{18}O_{V-SMOW}$ is due to the input of SiO₂-rich components rather than carbonate contamination.

Collectively, AFC processes can reproduce Os isotopic variations in the Tibetan ultrapotassic rocks but fail to fully recreate their Mg isotopic signatures. For the UPVR samples with low ¹⁸⁷Os/¹⁸⁸Os, their variably low δ^{26} Mg may be inherited from metasomatized mantle sources rather than obtained through contamination by varying crustal lithologies.

6.2. Geochemical fingerprints of mantle carbonatite metasomatism

6.2.1. Mantle metasomatism recorded in the entrained mantle xenoliths Metasomatized mantle sources are requisite for generating potassium-rich mafic rocks with enriched isotopic compositions and extremely high concentrations of incompatible elements (cf. Turner et al., 1996; Avanzinelli et al., 2009; Zhao et al., 2009; Liu et al., 2014b; Conticelli et al., 2015). For ultrapotassic magmatism in circum-Tethyan realm, the subduction of Tethyan seafloor has been invoked to supply oceanic sediments (i.e., terrigenous sediments and deep-sea carbonates) to refertilize mantle sources (e.g. Avanzinelli et al., 2009; Conticelli et al., 2015). Similarly, lithospheric mantle beneath the Lhasa terrane has been significantly metasomatized during the Tethyan seafloor subduction and the underthrusting of Indian continental lithosphere thereafter (cf. Chung et al., 2005; Zhao et al., 2009; Liu et al., 2014a).

Mineralogical and geochemical fingerprints preserved in mantle xenoliths provide important constraints on the geochemical evolution of mantle lithosphere. Recently, a high- δ^{18} O mantle region has been recognized beneath southern Tibet with the aid of insitu oxygen isotopic analyses on olivines separated from mantle xenoliths (Liu et al., 2014a). Accordingly, the primitive UPVR samples, with high MgO contents (up to 12.5 wt.%) and Mg[#] values (up to 76), exhibit oxygen isotopic compositions ($\delta^{18}O_{V-SMOW}$ = 7.6–8.1%, Zhao et al., 2009) that are comparable with this newly identified high- δ^{18} O mantle domain ($\delta^{18}O_{V-SMOW}$ up to 8.03%, Liu et al., 2014a). Besides, the low Ti/Eu (<1500) and high (La/Yb)_N (up to 89.5) features shown by clinopyroxenes with low Al_2O_3 contents in mantle xenoliths also suggest their origin of mantle carbonatite metasomatism (Coltorti et al., 1999) (Fig. S1). Considering that carbonatite metasomatism will increase clinopyroxene abundances in mantle assemblage at the expense of orthopyroxene (Coltorti et al., 1999; Huang et al., 2011), a high- δ^{18} O metasomatic agent like recycled marine carbonate (e.g. δ^{18} O for calcites in mantle peridotite is up to 20.5‰, Ducea et al., 2005) is needed to modify mantle lithology and oxygen isotopic composition beneath southern Tibet.



Fig. 5. Isotopic fingerprints of mantle carbonatite metasomatism recorded in the Tibetan ultrapotassic rocks. (a) $\delta^{26}Mg_{DSM3}$ vs. ⁸⁷Sr/⁸⁶Sr plot for less-contaminated ultrapotassic rocks with ¹⁸⁷Os/¹⁸⁸Os lower than 0.3 (i.e., UPVR samples in trend B, Fig. 3c), showing that both crustal contamination and mantle metasomatism have contributed to isotopic variations in the Tibetan UPVR. Carbonatitic metasomatism is modeled in terms of binary mixing between carbonates (Sr = 800 ppm, ⁸⁷Sr/⁸⁶Sr = 0.709) and mantle lherzolite (Sr = 21.1 ppm, ⁸⁷Sr/⁸⁶Sr = 0.703) (Table S7). AFC processes are modeled to show ⁸⁷Sr/⁸⁶Sr increments during crustal contamination. Varying ⁸⁷Sr/⁸⁶Sr and $\delta^{26}Mg$ are assumed for ultrapotassic magmas (Sr = 1200 ppm). AFC modeling curves are marked in 5% increments, where *R* is 0.8 and *D*_{Mg} is 4 ± 2. Bulk partition coefficient for Sr (*D*_{Sr}) is set at 0.7. Crustal contaminant is assumed to have radiogenic Sr isotopic composition (Sr = 320 ppm, ⁸⁷Sr/⁸⁶Sr = 0.756; Liu et al., 2014b), and its Mg isotopic compositin is similar to that of upper continental crust (Table S7). Shadowed area in Fig. 5a indicates crustal contamination occurred at relatively low level (AFC < 10%). Note that the variations in $\delta^{26}Mg$ are very limited when crustal contamination occurred at low level. (b) $\delta^{26}Mg_{DSM3}$ vs. Hf/Sm diagram, showing the dominant role of dolomites in changing mantle Mg isotopic composition. Given the intrinsic Mg isotopic variations in marine dolomites (Blättler et al., 2015) and their major and trace element changes during prograde metamorphism (Li et al., 2014), subducted dolomites, with ranges of $\delta^{26}Mg$, Hf/Sm, and MgO, are used for modeling carbonatitic metasomatism (see Table S7 for details). The binary mixing curves of mantle carbonatite metasomatism are marked in 10% increments and labeled with corresponding Mg/Ca ratio of recycled carbonates. The Hf/Sm and $\delta^{26}Mg$ data for oceanic basalts (MORB and OIB, Sun and McDonough, 1989; Teng e

6.2.2. Identifying mantle carbonatite metasomatism through Os–Sr–Mg isotopes

With respect to the metasomatic fingerprints left in mantle xenoliths, the relevance of carbonatite metasomatism and ultrapotassic magmatism in southern Tibet cannot be understated. Besides, potassium-rich silicate melts observed in melting experiments of carbonated pelite (850-1300 °C, 2.5-5.0 GPa, Thomsen and Schmidt, 2008) indicate the genetic links between carbonatite metasomatism and mantle-derived potassium-rich magmatism as well. If we considered that Lhasa terrane had drifted across the Tethys ocean during the Mesozoic (cf. Yin and Harrison, 2000; Zhu et al., 2013), the nature of subducted marine sediments thus directly affects the manner in which we interpret the geochemical signatures of the Tibetan ultrapotassic rocks. Previous works reveal that the Tethys ocean, which opened around the equator, may have greater carbonate compensation depth and accumulate more carbonates and organic-rich sediments due to the influence of equatorial upwelling (cf. Johnston et al., 2011). Moreover, the microbial dolomite model highlights the role of bacteria activity in overcoming kinetic barrier for dolomite nucleation at low-temperature, and promoting the precipitation of dolomite within anoxic, organic-rich deposits in deep-sea settings (cf. Mazzullo, 2000; Sánchez-Román et al., 2009; Blättler et al., 2015). The Mesozoic carbonate turbidite sequences found in Tethys domain (e.g. Wang et al., 2005) further indicate that shelf carbonates can be transported into or beyond the trench by turbidity current and finally involved in seafloor subduction (cf. Yang et al., 2012).

Combined with the extremely light Mg isotopic compositions in marine carbonates (Fantle and Higgins, 2014; Blättler et al., 2015), we suggest that carbonate-rich Tethyan sediments may have played a crucial role in replenishing mantle sources of the Tibetan ultrapotassic rocks. This suggestion is supported by the low Hf/Sm and high CaO/Al₂O₃ ratios in those primitive UPVR samples (Figs. 2c and 2d), because carbonatitic metasomatism will increase CaO contents of mantle sources and make mantle assemblages strongly depleted in HFSEs (e.g. low Hf/Sm) (Ducea et al., 2005; Huang et al., 2011). In addition, it is difficult for recycled marine carbonates, with low Os concentrations (Ravizza, 1993), to change mantle Os isotopic composition. Accordingly, carbonatitic metasomatism has potential to fully reproduce the considerable variation of δ^{26} Mg in those less contaminated ultrapotassic rocks (trend B in Fig. 3c). Recent study regarding Sr and Ca isotopes of Hawaiian shield lavas also highlights the presence of subducted carbonates in mantle sources and their capacity of raising 87 Sr/ 86 Sr in mantle lithology (Huang et al., 2011). Hence, mantle carbonatite metasomatism occurred beneath southern Tibet can also be manifested by the negative correlations between δ^{26} Mg and 87 Sr/ 86 Sr in the low- 187 Os/ 188 Os UPVR samples (Fig. 5a).

However, recycling marine carbonates into mantle sources cannot fully explain the variably high 87 Sr/86 Sr in ultrapotassic rocks (Fig. 5a), as the 87 Sr/ 86 Sr of subducted carbonates (≤ 0.71 , Fig. 5a) are not high enough to dramatically increase mantle Sr isotopic composition. Even if pronounced increases of ⁸⁷Sr/⁸⁶Sr can be observed when voluminous crustal materials were incorporated into ultrapotassic magma (>10% AFC, Fig. 5a), low-¹⁸⁷Os/¹⁸⁸Os signatures in these UPVR samples argue against extensive crustal contamination (Fig. 3a). Besides, much higher Sr concentrations in ultrapotassic magmas (most >1000 ppm, Table S2) relative to crustal contaminants will only give rise to minor increases in ⁸⁷Sr/⁸⁶Sr during low-level crustal contamination (shadowed area in Fig. 5a). Given that subducting sediments are overwhelmingly terrigenous (Plank and Langmuir, 1998), a considerable fraction of ⁸⁷Sr/⁸⁶Sr in the Tibetan UPVR could be due to the subduction of terrigenous sediments (GLOSS: 87 Sr/ 86 Sr = 0.717, Fig. 5a). In this scenario, the extremely radiogenic Sr isotopic compositions can be reconciled with the joint contribution from mantle metasomatism and low-level crustal contamination thereafter (cf. Liu et al., 2014b). It can be concluded that the variably lower δ^{26} Mg values, as well as negative Mg-Sr isotopic correlations defined by those less contaminated UPVR samples, provide evidence for the involvement of carbonate-bearing sediments in mantle refertilization beneath southern Tibet.

6.2.3. Implications for the effect of carbonatitic metasomatism on mantle Mg isotopic composition

Although both calcite and dolomite are reported to have low δ^{26} Mg values (Fantle and Higgins, 2014; Blättler et al., 2015), much more low-Mg carbonates are needed to modify mantle Mg isotopic

composition (Figs. 3c and 5a). Besides, compared with dolomiterich carbonates, calcite-rich carbonates cannot remain stable at high-pressure condition (Dasgupta and Hirschmann, 2010). Significant Mg isotopic exchanges between limestone and eclogitic oceanic crust further imply that calcite cannot keep its low δ^{26} Mg signature during prograde metamorphism (Wang et al., 2014). These findings cast doubts on the capacity of calcite-rich carbonates to cause Mg isotopic heterogeneity in mantle lithosphere.

For these reasons, we need a geochemical proxy for carbonatitic metasomatism to clarify which kind of marine carbonates can effectively change mantle Mg isotopic composition. Compared with ⁸⁷Sr/⁸⁶Sr and CaO, Hf/Sm ratio in mantle-derived rocks may be more suitable not only because carbonates are extremely depleted in Zr and Hf (Hf/Sm < 0.1, Fig. 2b), but because Hf and Sm have similar incompatibility during mantle melting and fractional crystallization (Sun and McDonough, 1989; Handley et al., 2011). The similar incompatibility between Hf and Sm during magmatism suggests that they will not fractionate from each other. Hence Hf/Sm variation in mantle-derived magmatic rocks can largely reflect the nature of mantle sources without being affected by different degree of mantle partial melting. This is consistent with the similar Hf/Sm ratios in ocean island basalts (Hf/Sm = 0.78) and mid-ocean ridge basalts (Hf/Sm = 0.78, Sun and Mc-Donough, 1989) (Fig. 5b). Additionally, the Hf/Sm of continental crust (0.68-1.13, Rudnick and Gao, 2003) and global subducting sediments (GLOSS, 0.70, Plank and Langmuir, 1998) are comparable with that of primitive mantle (0.70, Sun and McDonough, 1989), ruling out the interference from terrigenous sediment subduction and crustal contamination (Fig. 2b). Therefore, the combined use of δ^{26} Mg and Hf/Sm will allow us to better constrain mantle carbonatite metasomatism beneath southern Tibet.

Based on the intrinsic Mg isotopic variations in marine carbonates (Fantle and Higgins, 2014; Blättler et al., 2015) and their trace element variations during prograde metamorphism (Li et al., 2014), we modeled carbonatitic metasomatism in terms of binary mixing between mantle lherzolite and recycled carbonates with various Mg/Ca (Fig. 5b). It can be found that, with marine carbonates involved in the metasomatism, mantle peridotite becomes depleted in HFSEs and has lighter Mg isotopic composition (Fig. 5b). The poorly varying δ^{26} Mg indicate that recycled calcites cannot effectively change mantle Mg isotopic composition because their MgO contents are much lower than that of peridotite (Fig. 5b). Meanwhile, extensive isotopic exchanges with underlying eclogite during prograde metamorphism will even give rise to heavier Mg isotopes in calcite-rich carbonates (Wang et al., 2014), which makes it more difficult for recycled calcites to modify mantle Mg isotopic composition.

Conversely, subducted dolomites, with high MgO contents, are capable of preserving their initially light Mg isotopic compositions during prograde metamorphism (Wang et al., 2014), and thus have potential to leave Mg isotopic fingerprints in mantle lithosphere. As modeled in Fig. 5b, the δ^{26} Mg–Hf/Sm correlations in the Tibetan UPVR match well with the binary mixing trends defined by mantle lherzolite and recycled dolomites with Mg/Ca ranging from 1.04 to 0.60. Apart from the Tibetan ultrapotassic rocks, the potential of recycled dolomites to modify mantle Mg isotopic compositions is further revealed by δ^{26} Mg–Hf/Sm correlations in kamafugites from western Qinling orogenic belt, the Cretaceous basalts from north China craton (ca. 100 Ma, Yang et al., 2012), and the Cenozoic basalts from south China (13-3 Ma, Huang et al., 2015) (Fig. 5b). It seems that the co-variations between δ^{26} Mg and Hf/Sm in the UPVR not only provide strong argument for mantle carbonatite metasomatism beneath southern Tibet, but also illuminate the dominant role of recycled dolomite in modifying mantle Mg isotopic composition.



Fig. 6. Geochemical traverses (see Fig. 1a) showing spatial variation of Mg isotope and Hf/Sm ratio in ultrapotassic rocks. For comparison, two traverses have been corrected to positions relative to the LMF and SNMZ in the Lhasa terrane. Pink arrows indicate the enhancing role of mantle carbonatite metasomatism from south to north. The Hf/Sm range of arc magmas (White and Patchett, 1984) is shown for comparison.

6.3. Relevance of the Neo-Tethyan seafloor subduction and the generation of postcollisional ultrapotassic magmatism

Sedimentary contribution to mantle heterogeneity can be identified through trace element and isotopic geochemistry of mantlederived magmas (White and Patchett, 1984). Based on the discussion about Mg isotopic data and Hf/Sm variation in the UPVR, two geochemical traverses were carried out with the aim of further investigating mantle refertilization processes during the Tethyan seafloor subduction. As illustrated in Fig. 6, the UPVR samples in southern Lhasa subterrane display mantle-like Hf/Sm and δ^{26} Mg, but, in the central Lhasa subterrane, the Hf/Sm and δ^{26} Mg of the UPVR appear to be more variable. Due to the scarcity of UPVR outcropping in the northern Lhasa subterrane (Fig. 1a), it is untenable to ascribe the observed spatial variations of Mg isotope and trace element indicators to the southward subduction of Bangong-Nujiang Neo-Tethyan oceanic slab (cf. Zhu et al., 2013). We thus interpret these northward geochemical variations as representing a clear pattern of mantle refertilization during the northward subduction of the Indus-Yarlung Zangbo Neo-Tethyan oceanic slab. Because slab metamorphic dehydration would be gradually replaced by a melting regime at greater depth (Schmidt and Poli, 2003), the decreasing δ^{26} Mg and Hf/Sm from south to north may reflect the enhanced mantle carbonatite metasomatism with Neo-Tethyan oceanic slab going deeper (Figs. 6 and 7a). This interpretation is in agreement with the result of thermodynamic calculation (Connolly, 2005), which points out that significant amount of marine carbonates may have survived from the decarbonation and dissolution in sub-arc region and been transferred into deep earth.

In this regard, a two-stage petrogenetic model was proposed to link ultrapotassic magmatism in the Lhasa terrane and mantle refertilization during the northward subduction of Neo-Tethyan oceanic lithosphere. With the slab devolatilization processes gradually changed from metamorphic dehydration to melting regime at greater depths, partial melts of carbonate-bearing sediments



Fig. 7. A two-stage geodynamic model consists of (a) mantle carbonatite metasomatism during the northward subduction of the Neo-Tethyan seafloor, and (b) generation of the Oligocene–Miocene ultrapotassic magmatism in a tectonic setting of continental convergence. The devolatilization pattern during the northward subduction of the Neo-Tethyan oceanic lithosphere (>65 Ma) changed from metamorphic dehydration to a melting regime, which may correspond to more dolomites transported to the subcontinental lithospheric mantle. CC = Continental Crust; MBT = Main Boundary Thrust; MCT = Main Central Thrust; STDS = Southern Tibetan Detachment System.

stemmed from downgoing oceanic slab flux into the mantle wedge, leaving clinopyroxene-rich mantle assemblage (e.g. wehrlite, Yang et al., 2012) with isotopically light Mg (Fig. 7a). Under the background of continued underthrusting of Indian continent, thermal perturbation induced by deep geodynamic processes (e.g. convective thinning and/or delamination of the overthickened lithospheric mantle, roll-back and break-off of Indian slab) finally gives rise to the partial melting of metasomatized mantle sources and the occurrence of the Oligocene–Miocene ultrapotassic magmatism in southern Tibet (Fig. 7b). As evidenced by radiogenic Os isotopic compositions, entrained crustal xenoliths, and zircon xenocrysts with pre-eruptive U–Pb ages, ultrapotassic magmas underwent different extent of crustal contamination when they traveled through the overthickened Lhasa terrane crust.

7. Conclusions

Systematic analyses of Os, Sr, and Mg isotopes on the Tibetan ultrapotassic rocks elucidate the petrogenesis of potassium-rich mafic rocks and carbon recycling processes in orogenic belts. The variably high ¹⁸⁷Os/¹⁸⁸Os in mantle-derived ultrapotassic rocks demonstrate the widespread crustal contamination during magma ascent through the Lhasa terrane crust. But high MgO contents permit Mg isotopes in ultrapotassic rocks essentially unaffected by crustal contamination and enable Mg isotopic tracer to probe into the nature of mantle sources. The isotopically light Mg in those low-¹⁸⁷Os/¹⁸⁸Os ultrapotassic rocks is best interpreted to reflect the subduction of carbonate-bearing sediments into mantle sources, which are consistent with the negative Mg–Sr isotopic correlations and high CaO contents in those HFSE-depleted primitive ultrapotassic samples. The positive co-variations between δ^{26} Mg

and Hf/Sm further highlight the potential of recycled dolomite to change mantle Mg isotopic composition. The spatial variations in Hf/Sm and δ^{26} Mg correlate well with the expected changes in devolatilization regime during the northward subduction of Neo-Tethyan oceanic slab, suggesting that carbonatitic metasomatism operating in mantle refertilization beneath southern Tibet have laid the framework for generating ultrapotassic rocks in postcollisional stage.

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Appendix A. Supplementary material

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